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SHEET

1 OF 2

UNITED STATES PATENT DOCUMENTS				
EXAMINER'S INITIALS	CITE NO.	PATENT NUMBER	ISSUE DATE MM-DD-YYYY	FIRST NAMED INVENTOR
	A1			
	A2			
	A3			
	A4			

FOREIGN PATENT DOCUMENTS					
EXAMINER'S INITIALS	CITE NO.	DOCUMENT NUMBER	COUNTRY OR REGION	DATE OF PUBLICATION MM-DD-YYYY	FIRST NAMED INVENTOR OR APPLICANT
	B1				
	B2				
	В3				
	B4				

OTHER PRIOR ART - NON-PATENT DOCUMENTS				
EXAMINER'S CITE Include name of the author (in Capital Letters), title of the article (when appropriate), title of the item(book, journal, serial, symposium, catalog, etc.), date, page(s), volume-issue number(s), publisher, city and/or coupublished				
	C1	DUNN, C.J., WAGSTAFF, A.J., PERRY, C.M., PLOSKER G.L., GOA, K.L., Cyclosporin - An Updated Review of the Pharmacokinetic Properties, Clinical Efficacy and Tolerability of a Microemulsion-Based Formulation Neoral R(1) in Organ Transplanation, Drugs, 61: 1957-2016 (2001)		
	C2	PORTER, C.J.H., CHARMAN, W.N., In Vitro Assessment of Oral Lipid Based Formulations, Advanced Drug Delivery Reviews, 50: S127-S147 (2001)		
	C3	BALANDRAUD-PERI, N., OUENEAU, P.E., CAROLI-BOSC, F.X., BERTAULT-PERES, P., MONTET, A.M., DURAND, A., MONTET, J.C., Effects of Tauroursodeoxycholate Solutions on Cyclosporin and Bioavailability in Rats, Drug Metabolism and disposition, 25: 912-916 (1997)		

EXAMINER'S SIGNATURE	/Ivan Greene/	DATE CONSIDERED	04/03/2009

**EXAMINER:** Initial if reference considered, whether or not citation is in conformance with MPEP §609. Draw a line through citation if citation not in conformance and reference not considered. Include a copy of this form with next communication to applicant.

C4	GUO, J.X., PING, Q.N., CHEN, Y., Pharmacokinetic Behaviour of Cyclosporin A in Rabbits by Oral Administration of Lechithin Vesicle and Sandimmum Neoral, International Journal of Pharmaceutics, 216: 17-21 (2001)
C5	PORTER, C.J.H., CHARMAN, S.A., WILLIAMS, R.D., BAKALOVA, M.B., CHARMAN, W.N., Evaluation of Emulsifiable Glasses for the Oral Administration of the Cyclosporin in Beagle Dogs, International Journal of Pharmaceutics, 141: 227-237 (1996)
C6	LEIGH, M., HOOGEVEST, P.V., TIEMIESSEM, H., Optimising the Oral Bioavailability of the Poorly Water Soluble Drug Cyclosporin A Using Membrane Lipid Technology, Drug Delivery and Sciences, 1: 73-77 (2001)
C7	MIYAKE, K., ARIINA, H., IRIE, T., HIRAYMA, F., UEKAMA, K., Enhanced Absorption of Cyclosporin A by Complexation with Dimethyl-Beta-cyclodextrin in Bile duct-cannulated and Non- Cannulated Rats, Biological and Pharmace
C8	BONDUELLE, S., CARRIER, M., PIMIENTA, C., BENOIT, J.P., LENAERTS, B., Tissue Concentration of Nanoencapsulated Radiolabelled Cyclosporin Following Peroral Delivery in Mice or Ophthalmac Application ni Rabbits, European Journal of Pharmaceutics and Biopharmaceutics, 42: 31-319

EXAMINER'S	/Ivan Greene/	DATE	04/03/2009
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**EXAMINER:** Initial if reference considered, whether or not citation is in conformance with MPEP §809. Draw a line through citation il citation not in conformance and reference not considered. Include a copy of this form with next communication to applicant.